Photobiomodulation for Hypertension and Alzheimer’s Disease

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Abstract. Although the cause(s) of Alzheimer’s disease in the majority of cases remains elusive, it has long been associated with hypertension. In animal models of the disease, hypertension has been shown to exacerbate Alzheimer-like pathology and behavior, while in humans, hypertension during mid-life increases the risk of developing the disease later in life. Unfortunately, once individuals are diagnosed with the disease, there are few therapeutic options available. There is neither an effective symptomatic treatment, one that treats the debilitating cognitive and memory deficits, nor, more importantly, a neuroprotective treatment, one that stops the relentless progression of the pathology. Further, there is no specific preventative treatment that offsets the onset of the disease. A key factor or clue in this quest for an effective preventative and therapeutic treatment may lie in the contribution of hypertension to the disease. In this review, we explore the idea that photobiomodulation, the application of specific wavelengths of light onto body tissues, can reduce the neuropathology and behavioral deficits in Alzheimer’s disease by controlling hypertension. We suggest that treatment with photobiomodulation can be an effective preventative and therapeutic option for this neurodegenerative disease.

Keywords: Cell death, infrared, mitochondria, non-pharmacological, red, vascular pathology

INTRODUCTION

Alzheimer’s disease (AD) is a brain disorder that involves a progressive loss of memory, cognition, and self-control, the so-called executive brain functions. It affects mainly those entering older age (>65 years); in fact, age is the major risk factor of the disease [1, 2]. The devastating symptoms arise after a large number of neurons across the brain degenerate. These dying neurons have a number of distinct features (Fig. 1). Intracellularly, neurons develop neurofibrillary tangles, made up mainly of hyperphosphorylated tau protein, that disrupt normal cell function and survival. There are also problems with glucose metabolism that will eventually lead to damage and dysfunction of the “engine room” of the cells, namely the mitochondria [3]; further, there are deficits evident of the intracellular clearance system, for example of proteins and glucosconjugates that involve small organelles called lysosomes. In addition to these intracellular features, there are also many extracellular structures, made up of aggregations or plaques of amyloid-β (Aβ) protein, scattered across the brain, particularly within the memory areas [4–7]. Finally, there is evidence of vascular pathology. In particular, there are sites of micro-bleeding across the brain, many of which are associated with the Aβ plaques [8–12].

In the sections that follow, we will examine briefly the suspected causes and mechanisms that underpin AD, focusing on the idea that hypertension and dysfunction of the vascular system may be central to the disease process. Next, we consider the current treatment options and preventative measures available for...
patients. We will then introduce the method of photobiomodulation, the use of red to near infrared light ($\lambda = 600$–$1300$ nm) on body tissues, and consider its beneficial effects on cells and on the vascular system and hypertension [13]. Finally, we will propose that photobiomodulation can form an effective preventative and therapeutic treatment option for patients by controlling hypertension.

THE CAUSES AND MECHANISMS: A FOCUS ON THE VASCULAR SYSTEM

Over the years, many hypotheses as to the causes and mechanisms underpinning the pathology associated with AD have been proposed and debated [1–7, 14]. Briefly, these include suggestions that the primary driver of the neuronal dysfunction and death, either by genetic mutation or by factors unknown, is either the accumulation of Aβ, the formation and self-propagation of tau protein and/or the dysfunction of mitochondria and/or lysosomes. Unfortunately, each one of these hypotheses has been associated with problems, in particular, in the development of a curative therapy [15, 16], with none being able to fully account for all the features of the disease [4–7].

Vascular pathology and hypertension

One of the first hypotheses of the initial causes of dementia and AD, and a major focus of this review, centers on vascular pathology and cerebral hypoperfusion [17, 18]. This process may trigger the development of Aβ plaques, tau assemblies, mitochondrial and lysosome dysfunction, and ultimately neuronal death. There is evidence of cerebral micro-bleeding across the capillary-bed, together with damage to the capillary endothelium and blood-brain barrier [9, 19–21]. Further, from postmortem material, a majority of patients diagnosed with AD (90%) have been found to have considerable vascular pathology [18].

Hypertension has long been considered to be a major cause of this vascular pathology; it is also considered to be a strong risk factor for the development of AD [22–24]. Many studies have reported that systolic hypertension, as well as overall variability in blood pressure, is associated with memory impairment and development of AD [22, 25–30]. This increase in blood pressure leads to damage to the cerebral microcirculation and arterial stiffness [31], inducing ischemia, disruptions to the blood-brain barrier, loss of neuronal function and an increase in Aβ expression [32–35]. There are many fac-
tors that may contribute to hypertension, including alterations of the renin–angiotensin–aldosterone system or autonomic nervous system, an increase in salt intake, obesity, and insulin resistance [36–38]. The renin–angiotensin–aldosterone system is a key regulator of blood volume and systemic vascular resistance. Unlike the baroreceptor reflex that responds to acute changes in blood pressure, the renin–angiotensin–aldosterone generates more chronic alterations. The major compounds, namely renin, angiotensin II, and aldosterone, work together to raise blood pressure in response to various factors, including low blood pressure and/or salt levels [39–41]. However, it should be noted that, as with most hypotheses associated with neurodegenerative disease, the link between cognitive decline and hypertension has been subject to some debate over the years. While many studies have indicated increased cognitive impairment risk with hypertension (see above), other studies have suggested less of an association [26, 42, 43]. The timeline of hypertension may explain this discrepancy; in general, subjects with chronic hypertension over a period greater than ten years appear to have a stronger risk of cognitive decline later in life [44, 45].

Further to this debate, many clinical studies have reported that medications aimed to reduce hypertension lead to a lower incidence of dementia and AD [46, 47]. A recent meta-analysis containing fourteen international studies have highlighted considerable cognitive enhancement with blood-brain barrier-crossing renin-angiotensin drugs [48]. In addition, there is evidence in transgenic mouse models of the disease (APP/PS1, 3xTgAD) that hypertension can exacerbate the behavioral deficits and neuropathology of AD. For example, hypertensive APP/PS1 mice develop an earlier cognitive deficit, together with an increase in both Aβ plaques and levels of soluble Aβ in the brain and plasma when compared to normotenive APP/PS1 mice. The hypertensive APP/PS1 mice also have considerably more vascular pathology (e.g., reduced density of microvessels) than the normotenive APP/PS1 mice [49]. In the 3xTgAD mouse model, similar patterns have been reported, with experimentally-induced hypertension triggering an acceleration of Alzheimer-like pathology, microglial activation and leakage of the blood-brain barrier [50]. In addition, several authors have tested the efficacy of various drugs, in particular RB150 [51], in controlling hypertension in the 3xTgAD mouse model, with some promising outcomes [52].

It should be noted that vascular pathology and hypertension have been found in other types of neurodegenerative disease, not just AD. For example, in patients with Parkinson’s disease, there are reports of damage to the capillary endothelium and blood-brain barrier in the basal ganglia together with some fluctuation in blood pressure, including nocturnal hypertension [53, 54].

Taken all together, hypertension can lead to vascular pathology, altering the structure and density of cerebral blood vessels. This process can be disruptive to the mechanisms that assure an adequate blood supply to the brain, leading to cerebral hypoperfusion and potentially dementia and AD [22].

CURRENT TREATMENT OPTIONS

There are few effective treatment options available for patients with AD [55]. Some patients appear to respond to a newly approved therapy that aims to remove the Aβ plaques from the brain, while others gain some benefit with the use of acetylcholinesterase inhibitors, that suppress the breakdown of acetylcholine, and N-methyl-D-aspartate receptor antagonists, that helps prevent neurotoxicity in the brain [15, 56]. However, a number of patients do not respond to these treatments, and many suffer severe side effects. In essence, there is no currently available, broad range treatment option that stops or even slows the neuronal death (i.e., a neuroprotective treatment), although several experimental therapies have been suggested [57, 58]. The disease progresses relentlessly, and neurons continue to degenerate across the brain. Further, there is no treatment that even treats effectively the symptoms of the disease, namely the decline in memory, cognition, and self-control [59]. Indeed, the greatest fear for most people entering older age is that they will be diagnosed with AD and that there would be nothing that can be done about it.

In terms of prevention, a number of measures, albeit largely not specific for AD, have been suggested over the years, including, to do more exercise, be cognitively active, eat a Mediterranean diet, reduce both alcohol intake and obesity, and stop smoking [1, 7]. These measures can also help maintain a healthy blood pressure, controlling hypertension. In addition, and relevant to this review, suggestions have been made, based on evidence from both humans and animal models, that controlling hypertension can be an effective means to offset the onset of the disease,
together with limiting its severity (see above). Hence, a key to a better and more efficient treatment for AD may lie in addressing the control of hypertension.

PHOTOBIOMODULATION: THE LIGHT

In this context, there is a new treatment option that has been receiving considerable interest across the clinical, scientific, and wider community. This treatment is known as photobiomodulation, the use of red to near infrared light (\(\lambda \approx 600–1300\) nm) on body tissues [60]. It has been shown to be beneficial in many animal models of disease, from Parkinson’s disease [61] to stroke [62] and from traumatic brain injury [63] to retinal disease [64] and multiple sclerosis [65], it has an impeccable safety record, with little or no evidence of side effects or toxicity on body cells and the devices are easy to use and inexpensive [60]. In the sections that follow, the mechanisms of photobiomodulation are considered, together with its impact on the vascular system, hypertension, and AD.

The light mechanisms: direct and indirect stimulation

The precise mechanisms used by photobiomodulation to achieve beneficial outcomes are not entirely clear, but two have been suggested, namely direct and indirect stimulation [66, 67]. For direct stimulation, photobiomodulation has to fall directly on the neurons (Fig. 2). This stimulates the activity of the mitochondria, by being absorbed by a photosceptor (e.g., cytochrome oxidase c and/or interfacial water), whose primary function is to provide energy for the cells that drive many intrinsic functions. This process leads to a healthier cell, with the expression of various functional and protective genes that make the cell not only function better, but make it more resistant to distress and disease [60]. Many studies have also reported that photobiomodulation promotes many other beneficial outcomes, such as triggering neurogenesis and synaptogenesis, as well as reducing inflammation [60, 68–72].

In addition to direct stimulation, photobiomodulation has been shown, quite remarkably, to be beneficial to neuronal function and survival even when it is applied to a more distant or remote location; that is, when it is not applied directly to the neurons (Fig. 3). This indirect stimulation is thought to activate circulating cells or molecules, or even free-floating mitochondria, within the cardiovascular or lymphatic systems that then leads to an increase in overall mitochondrial activity, in a similar fashion to the direct stimulation described above, in the distressed neurons located in the brain. The indirect stimulation, although offering neuroprotection, is less effective than direct application [66, 67].

Lights on the vascular system and hypertension

Photobiomodulation has also been shown to have a direct effect on the vascular system. There is evidence that photobiomodulation offsets the degeneration and leakage of retinal capillaries in animal models of diabetes [73] and in the striatum and brainstem of
a mouse model of Parkinson’s disease [74]. Furthermore, photobiomodulation has been reported to induce the release of nitric oxide from cells, which triggers the vasodilation of nearby blood vessels, increasing blood (and lymphatic) flow [60, 75]. In addition to these studies, several groups have indicated an effect of photobiomodulation on the vascular system and cerebral blood flow. Photobiomodulation has been shown to improve endothelial dysfunction [76], influence the levels of angiogenic protein (e.g., vascular endothelial growth factor), that prompts capillary-like structure formation [63, 68], together with cerebral hemodynamic and uric acid levels [77], both of which are risk factors for the development of hypertension. There are also indications that photobiomodulation can increase oxygenated hemoglobin and oxidized cytochrome c oxidase levels [78], as well as restoring capillary blood supply and stimulating the microcirculation in Parkinson’s disease [79]. In AD patients, and in a patient suffering persistent vegetative state, photobiomodulation has been reported to increase cerebral blood flow [80–82]. In patients with major depressive disorder, photobiomodulation induces an improvement in diastolic blood pressure and a better cerebral blood flow, together with reductions in anxiety and depressive scores [83, 84].

There is also recent evidence in various experimental animal models that photobiomodulation reduces hypertension. In one model, involving constriction of the renal artery, photobiomodulation generates a long-lasting hypotensive effect after a device was implanted within the abdominal aorta. This effect is thought to be prompted by a photobiomodulation-induced vasodilation after a release of nitric oxide [85]. In another model of hypertension, involving an implantation of a cannula into the femoral artery, photobiomodulation applied externally over the abdomen reduced blood pressure considerably, together increasing serum levels of nitric oxide [86, 87].

**Lights on Alzheimer’s disease**

There are many studies, mainly on animal models and but also on patients, reporting that photobiomodulation is effective in AD. In animal models, that include different types of transgenic or Aβ-induced models, photobiomodulation has been shown to reduce Aβ plaques and aggregations, neurofibrillary tangles of hyperphosphorylated tau protein, synaptic vulnerability, gliosis, inflammation and oxidative stress, and increase in neurogenesis, mitochondrial function, and ATP levels (APP/PS1 model [88–93]; TASTPM model [94]; K3691 model [89, 90]; Tg2576 model [95]; Aβ-induced model [96–98]; 5xTg model [99]; hTau and 3xTg models [100]). In addition, after internalization of Aβ into human neuroblastoma cells, photobiomodulation has been shown to increase ATP levels and cell number, while reducing Aβ aggregates [101]. When applied to normal mice, photobiomodulation prompts a much faster clearance of Aβ out of the brain and into the lymphatic system [75]. Finally, there are also some very early, encouraging observations in AD patients that photobiomodulation improves cognition [81, 102–104] and the functional connectivity of different areas of the brain [82].

From these previous studies, there are two key and relevant issues worthy of further comment. First, in both the rodent models and patients, photobiomodulation was applied transcranially. Many previous studies have reported that light can penetrate up to 30–50 mm of body tissue; in general, the longer the wavelength then the greater the penetration [60, 67, 105, 106]. In rodents, where the distances between the scalp and the most ventral areas of the brain are relatively short (5–10 mm), photobiomodulation applied in this way can reach and influence all regions of the brain directly. In addition, transcranial photobiomodulation can reach the many cerebral and scalp blood and lymphatic vessels and hence influence circulatory mitochondria, cells, and molecules. In humans, where the distances are much greater (80–100 mm), transcranial photobiomodulation can still reach the superficial areas of the cerebrum where much of the AD pathology is located, namely the cortex (10–15 mm) [66, 67]. As with the rodents, transcranial photobiomodulation in humans can also reach the cerebral and scalp blood and lymphatic vasculature. Hence, in both rodents and humans, transcranial photobiomodulation, by reaching large areas of brain together with blood and lymph vessels, can impart beneficial outcomes by both direct and indirect stimulation (see above). It should be noted that in one study in mice, beneficial outcomes were achieved after photobiomodulation was applied across the abdomen only, indicating that indirect stimulation by itself can be effective in AD [96].

Second, in all of these previous studies on animal models, photobiomodulation was started either at the onset, or well after the onset, of the first behavioral deficits and/or pathology [88–100]. This approach is largely in line with the clinical reality...
of the disease, where patients are treated after the emergence of the first symptoms. However, for a preventative approach, it would be more instructive to treat animals well before the onset of pathology and behavioral signs. In one study, using a mouse model of ageing (CD1), a preventative approach was used and improvements were noted in working memory [107]. Although these findings are encouraging, this was not a model of AD.

PHOTOBIO MODULATION, HYPERTENSION, AND ALZHEIMER’S DISEASE: AN HYPOTHESIS AND A POSSIBLE TREATMENT APPROACH

When considering all this information, principally from animal models, we would like to highlight three key issues, that:

1. photobiomodulation reduces the pathology and behavioral deficits in AD when applied directly onto the head or body;
2. photobiomodulation can reduce hypertension and vascular pathology when applied to blood vessels, and;
3. hypertension exacerbates the neuropathology and behavioral deficits of AD.

Taking these three key issues together, we hypothesize that photobiomodulation, by controlling hypertension, can be an effective therapy option for AD. If applied at very early stages, photobiomodulation may form a preventative therapy by helping to maintain low blood pressure, thereby offsetting the onset of vascular pathology and the disease process; if applied at later stages, it may help form an ongoing therapeutic treatment, by controlling the hypertension, thereby slowing the disease process.

One of the clear advantages for using photobiomodulation is that, unlike many pharmaceutical options, there are few or no side effects, no evidence for toxicity on body cells and because the devices are easy to use and inexpensive, many patients can afford the treatment and use it at home. In fact, it has been our experience with six Parkinson’s disease patients that compliance to using photobiomodulation is very high [108]; a major factor in this high compliance rate was that photobiomodulation is a non-pharmacological treatment and that the patients felt at ease with its use.

In terms of approach, we suggest that photobiomodulation could be applied transcranially, hence offering both mechanisms of direct and indirect stimulation (see above); in addition, using this approach, the access to both cerebral and scalp blood vessels would provide photobiomodulation the potential to influence hypertension. One could also have the additional approach of applying photobiomodulation across another part of the body, for example the abdomen, as to access more blood vessels and influence hypertension. There is evidence, particularly using the longer wavelengths, that transabdominal photobiomodulation can pierce through to the intestines and influence composition of the microbiome in humans [109]; hence, if it can penetrate through layers of abdominal fat, muscle, and connective tissue, it would certainly reach the rich array of major blood vessels that innervate the abdominal organs (e.g., aorta). It could also influence the composition of the microbiome by reducing systemic inflammation. As a starting point, we suggest using a transcranial helmet [82, 103, 108] and a transabdominal device [109] set at the parameters used by previous studies. There is also the option of using a whole body photobiomodulation bed.

CONCLUSIONS

There is a real need for an effective, broad range therapeutic treatment option for patients suffering AD, one that treats the symptoms and offers neuroprotection. Further, is a need for a preventative treatment, one that can offset the development of the disease later in life. In this context, many previous studies have reported that photobiomodulation reduces the pathology and behavioral deficits in animal models of AD; there are also reports that photobiomodulation can reduce hypertension and limit the vascular pathology, a factor that has been reported to increase the risk of developing the disease. When taking all these findings together, we suggest that photobiomodulation, by controlling hypertension, could form an effective therapy for AD. We propose that early treatment may help maintain low blood pressure and offset the vascular pathology and development of the disease (preventative treatment), while later treatment may help control hypertension and slow the ongoing disease process (therapeutic treatment). The stage is set for a series of pre-clinical experiments using a combined hypertensive-AD animal model, translating to a large-scale clinical trial in humans.
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REFERENCES


erates the progression of Alzheimer-like pathology in a mouse model of the disease. \textit{Hypertension} 65, 218-224.


